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- (54) Gene encoding enzyme having flavin reducing activity and nitroreductase activity.
- The present invention succeeds in isolating a gene encoding an enzyme having an FMN reducing activity and a nitroreductase activity derived from luminous bacteria Vibrio fischeri (ATCC 7744), elucidating its primary structure, and producing Escherichia coli which can express the gene in large quantities. That is, the present invention provides a gene encoding an enzyme having the flavin reducing activity and the nitro-reductase activity, an enzyme produced therefrom, a recombinant vector containing the gene, and bacteria containing the recombinant vector.

Fig. 1	
ATC ACL CAR CCL ATE ATE CAR CAR ATE GAT ANY EGG TAR ACL QES AAJ Wet The Dis Pro lie lie His Asp Leu Giu Ash ark Try The See Lys	48
1 5 15 AAJ TAK GAR CCL ORS AAJ AAJ GTE ORS CAJ GAJ GAR ATY GCE GTE ATY	96
Lys Tyr Asp Pro Ser Lys Lys Val Ser Glu Glu Asp Leu Ala Val Leu	B10
20 25 30	
ATY GAJ GUL ATY FOZ ATY QES CUL ORS ORS ATM AAR ORS CAJ CUL TGG	144
Leu Glu Alu Leu Arg Leu Ser Aln Ser Sei 11e Asa Ser Glu Pro Trp 35 40 45	
AAJ TIK ATW UTL ATW GAJ ORS GAK GCA GCL AAJ CAJ GGL ATG CAK GAK	192
Lys Phc He Val He Glu Ser Asp Aln Ata Lys Glu Gly Wet His Asp	
50 55 60	
ORS TIK GOL AAR ATG CAK CAJ TIK AAK CAJ COL CAR ATR AAT GOL TGR	248
Ser Phe Alu Asn Mei Ris Glo Phe Asn Glo Pro His He Lya Ata Cys 65 70 75 80	
ORS CAR GTG ATH ATY THE GGL AAR AAJ ATY ORS TAK ACL WOX GAR GAR	288
Ser His Val 11c Lea Phy Ala Asa Lya Lea Ser Tyr Thr Arg Aup Asp	
85 90 95	
TAK CAK CTC GTL KTY QRS AAJ CCL CTL GCL CAK AAJ 16Z ATW ACL GAJ Tyr Asp Yul Yal Leu Ser Lys Ata Yul Ala Asp Lyu Arg 11c Thy Ciu	236
100 105 110 All 105	
GAU CAJ AAJ GAJ GCL CCL TIK GCL QRS TIK AAJ TIK CIL CAJ TIG AAK	384
Glu Glu Lys Glu Ala Ala Phe Ala Ser Phe Lys Phe Val Glu Len Asa	
115 120 125	
THE GAK GAJ AAK GOL GAJ CAK AAJ COL THE ACL AAJ COL CAI GOL TAK Cys Asp Glu Ash Cly Glu His Lys Alb Tep The Lys Pro Glu Alb Tye	432
130 135 140	
NTY GCL NTY CGL AAR GCL NTY CAN ALL NTY GCL PGZ NTY AAR ATH GAN	480
Les Ala Les Gly Asn Ala Les His Thr Les Ala Arg Les Asn 11c Asp	
145 150 155 160	
ORS ACL ACL ATC GAJ GGL ATM GAK CCL GAJ ITY TTG ORS GAJ ATM TTK Ser Thr Thr Net Gin Gly He Asp Pro Gla Leu Leu Ser Glu He Phe	528
160 170 175	
OCL GAN GAJ RTY AAJ OUR TAK DAJ TOK CAR GTE GOL ATY GOL ATH GOL	57G
Alu Asp Glu Leu Lys Gly Tyr Glu Cys His Yal Ala Leu Ala lle Gly	
180 185 190	
TAN CAN CAN COL ORS GAT GAN TAN AAN GOL ORS THE COL AND ORS TOZ THE RES RES Pro See Gio And The Ann Ain See Lea Pro Lys See Arn	624
195 200 205	
AAJ GCL TTR GAJ GCL GTL ATM ACL ATM ATY TJJ	651
Lys Ala Phe Giu Ala fai lie Thr lie Leu 444	
210 215	

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TITLE OF THE INVENTION

Gene Encoding Enzyme Having Flavin Reducing Activity and Nitroreductase Activity

BACKGROUND OF THE INVENTION

(i) Field of the Invention

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The present invention relates to a gene encoding an enzyme having a flavin reducing activity and a nitroreductase activity, the enzyme produced therefrom, a recombinant vector containing the gene and bacteria containing the recombinant vector.

(ii) Description of the Related Art

A bacterial luciferase derived from luminous bacteria is used to produce oxidized flavin adenine mononucleotide (hereinafter referred to as "oxidized FMN") and a long-chain carboxylic acid in the presence of a long-chain aliphatic aldehyde, oxygen and a reduced flavin adenine mononucleotide (hereinafter referred to as "FMNH₂") as a luminescent substrate, and in this case, the bacterial luciferase catalyzes a reaction in which blue light is emitted. FMNH₂ which is a substrate can be obtained from a reduced nicotinamide adenine dinucleotide:flavin mononucleotide (NADH:FMN) reductase and a reduced nicotinamide adenine dinucleotide phosphate:flavin mononucleotide (NADPH:FMN) reductase, and the long-chain aldehyde can be obtained from a fatty acid reductase complex.

In recent years, Spyrou et al. have isolated a flavin reductase gene from Escherichia coli and elucidated its primary structure, which is disclosed in Spyrou G., Haggard-Ljungquist E., Krook M., Jornvall H., Nilsson E. and Reichard P., J. Bacteriol, 173, p. 3673-3679 (1991).

Around us, there are many substances (mutagens) which damage chromosomal DNA, and during our lives we are exposed to these substances. Nitroarenes are members of one group of environmental mutagens, and they are contained in the exhaust gas of automobiles, the smoke of incinerators, the atmosphere of cities, the bottoms of rivers, the air in rooms where stoves are lighted, and the burnt portions of grilled chickens. Of nitroarenes having mutability and carcinogenicity, 2-nitrofluorene is well known.

A nitroarene itself does not react directly with DNA to damage the same, but a metabolite of the nitroarene gives rise to a mutation in DNA to damage the DNA. For example, it can be presumed that nitrofluorene is reduced to an N-hydroxy form in the cell of a microorganism by a nitroreductase and then activated by an o-acetyl transferase, to thus finally produce nitrenium ions which attack the DNA. Therefore, it can be considered that the reaction of the nitroreductase with 2-nitrofluorene is a rate determining step in the mutagenesis of DNA by 2-nitrofluorene.

Watanabe et al. have isolated a nitroreductase gene from <u>Salmonella</u>, which is disclosed in Watanabe M., Ishidate M, Jr and Nohmi T., Mutation Research, p. 216 211-220 (1989). Furthermore, its primary structure has been elucidated in Watanabe M., Ishidate M, Jr and Nohmi T., Nucleic Acid Research, <u>18</u>, p. 1059 (1990).

As understood from the foregoing, the FMN reductase is essential to utilize the luminescent reaction of bacterial luciferase to the utmost. Therefore, the isolation of the FMN reductase gene permits preparing the enzyme in large quantities, and thus, an important object is the isolation of the gene encoding this enzyme.

Furthermore, the nitroreductase gene is useful to improve the detection sensitivity of the above-mentioned mutagen or carcinogen.

However, with regard to the isolation of the FMN reductase gene of luminous bacteria and the nitroreductase gene as well as the expression of them in <u>Escherichia</u> <u>coll</u>, no report has been made so far.

SUMMARY OF THE INVENTION

In view of the above-mentioned technical situation, an object of the present invention is to provide a gene encoding an enzyme having an FMN reducing activity of luminous bacteria and a nitroreductase activity and the enzyme therefor. Another object of the present invention is to provide a recombinant vector containing this gene and bacteria containing the recombinant vector.

As a result of intensive research, the present inventors have succeeded in isolating a gene encoding an enzyme having the FMN reducing activity and the nitroreductase activity from the luminous bacteria <u>Vibrio fischeri</u> (ATCC 7744), and in elucidating its primary structure. In addition, they have succeeded in cultivating <u>Escherichia coli</u> transformed with a vector containing the gene which can express the protein in large quantities. As a result, the present invention has now been completed.

The present invention has the following parts (1) to (8).

- (1) A gene containing a nucleotide sequence shown in Fig. 1 and encoding an enzyme having a flavin reducing activity and a nitroreductase activity.
- (2) A gen containing a nucleotide sequence shown in Fig. 2 and encoding an enzyme having the flavin reducing activity and the nitroreductase activity described in the previous paragraph (1).
- (3) A gene containing a nucleotide sequence shown in Fig. 3 and encoding an nzyme having a flavin reducing activity and a nitroreductase activity.
- (4) An enzyme containing an amino acid sequence shown in Fig. 4 and having a flavin reducing activity and a nitroreductase activity.
- (5) A recombinant vector containing a DNA whose nucleotide sequence is shown in Fig. 1.
- (6) The recombinant vector described in the previous paragraph (5) in which the gene having the nucleotide sequence shown in Fig. 1 is inserted into a plasmid vector.
- (7) Bacteria containing a recombinant vector containing a DNA whose nucleotide sequence is shown in Fig. 1.
- (8) A method for preparing an enzyme containing an amino acid sequence shown in Fig. 4 which compris s the step of cultivating bacteria transformed with a recombinant vector containing a DNA whose nucleotide sequence is shown in Fig. 1.

BRIEF DESCRIPTION OF THE DRAWINGS

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Fig. 1 shows a nucleotide sequence of a gene encoding an enzyme having a flavin reducing activity and a nitroreductase activity.

Sequence length: 657

Sequence type: Nucleic acid

Strandedness: 1
Topology: Linear

Molecular type: Genomic DNA Feature of sequence description:

Feature key defined in Gene Bank Authorin

30 Reference Manual Release 1.1 (hereinafter referred to simply as "feature key"): CDS

Procedure for determining the feature:

Prediction from an amino acid sequence of Fig. 4 based on genetic code table.

(On the left and right sides of each triplet, a 5' terminal and a 3' terminal are present, respectively. This triplet represents a purine base (Pu) and a pyrimidine base (Py) constituting a nucleotide sequence.

- 35 A: adenine,
 - G: guanine,
 - C: cytosine,
 - J: A or G,
 - K: Tor C.
- 40 L: A, T, C or G,
 - M: A, C or T,
 - T: thymine,
 - X: when Y is A or G, X is T or C, or when Y is C or T, X is C,
 - Y: when X is C, Y is A, G, C or T, or when X is T, Y is A or G,
 - W: when Z is C or T, W is C or A, or when Z is C or T, W is C,
 - Z: when W is G, Z is A, G, C or T, or when W is A, W is A or G,
 - : when S is A, G, C or T, QR is TC, and *** represents TAA, TAG or TGA.).

Under each triplet codon of the nucleotide sequence, the amino acid encoded thereby is represented.

Fig. 2 shows a typical nucleotide sequence of the gene encoding the enzyme having a flavin reducing activity and a nitroreductase activity.

Sequence length: 657

Sequence type: Nucleic acid

Strandedness: 1 Topology: Linear

55 Molecular type: Genomic DNA

Original source:

Organism: Vibrio fischeri Strain: ATCC 7744

Feature of sequ nce description:

Feature k y d fined in Gene Bank Authorin Referenc Manual R lease 1.1: CDS

Procedure for determining the f ature:

Experimental procedure.

5 Fig. 3 shows a nucleotide sequence of a gene encoding an enzyme having a flavin reducing activity and a nitroreductase activity.

> Sequence length: 929 Sequence type: Nucleic acid

Strandedness: 1 Topology: Linear

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Molecular type: Genomic DNA

Original source:

Organism: Vibrio fischeri Strain: ATCC 7744

Feature of sequence description:

Feature key: CDS

Site having the feature: 109-762

Procedure for determining the feature:

Experimental procedure.

20 Fig. 4 shows an amino acid sequence of an enzyme having a flavin reducing activity and a nitroreductase activity.

Sequence length: 218
Sequence type: Amino acid
Molecular type: Protein.

Fig. 5 shows an N-terminal amino acid sequence of an NAD(P)H:FMN reductase and synthetic oligonucleotide probes (FR1 and FR2).

Fig. 6 shows a restriction map of the gene of the present invention and a sequencing strategy. Arrows denote directions for the determination of the nucleotide sequences. The portion indicated by a box corresponds to the gene.

Fig. 7 shows a process of constructing a recombinant vector (an expression vector pFR7) containing a gene encoding an enzyme having an FNM reducing activity and a nitroreductase activity of luminous bacteria according to the present invention.

Fig. 8 shows the confirmation of the expressed protein by SDS-polyacrylamide gel electrophoresis. Lane 1 is a pUC8/D1210 strain, lane 2 is a pFR7/D1210 strain, lane 3 is a pFR5/D1210 strain, and lane 4 is a Boehringer Mannheim NAD(P)H:FMN reductase.

The symbols used in the drawings have the following meanings.

lacP lactose promoter

Ampr Ampicillin resistant gene

pUC8 plasmid vector

pFR3 recombinant vector

pFR5 recombinant vector

pFR7 recombinant vector (expression vector)

Sau3Al four bases (GATC) recognizing restriction enzyme
HincII six bases (GTPyPuAC) recognizing restriction enzyme

Smal six bases (CCCGGG) recognizing restriction enzyme

Stul six bases (AGGCCT) recognizing restriction enzyme

EcoRI six bases (GAATTC) recognizing restriction enzyme

HindIII six bases (AAGCTT) recognizing restriction enzyme

Accl six bases (GTATAC) recognizing restriction enzyme

50 NspV six bases (TTCGAA) recognizing restriction enzyme

EcoRV six bases (GATATC) recognizing restriction enzyme.

DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS

The gene of the present invention is characterized by containing a nucleotide sequence having a sequence I ngth of 657 bases as shown in Fig. 1. The nucleotide sequence in Fig.1 can be predicted from an amino acid s quence shown in Fig. 4 as mentioned later.

A preferable sequence contains a nucleotide sequence as shown in Fig. 2.

A typical nucleotide sequence is a DNA having a sequence length of 929 bases as shown in Fig. 3.

Th basic nucleotides quence of the present invention is deriv d from Genomic DNA isolated from luminous bacteria Vibrio fischeri (ATCC 7744). This sequence is characterized by encoding a protein having a molecular weight of 24562 and comprising 218 amino acids, corresponding to nucleotides numb red 109 to 762.

The gene of the present invintion encodes a protein having a flavin reducing activity and a nitroreductas activity, for example, an FMN reducing activity and a nitrofurazone reducing activity.

An enzyme of the present invention is a protein having an amino acid sequence shown in Fig. 4 which can be predicted from the nucleotide sequence in Fig. 3. This protein comprises 218 amino acids and has a molecular weight of 24562 and the two activities of luminous bacteria, i.e., the flavin reducing activity and the nitroreductase activity.

A recombinant vector of the present invention contains a DNA whose nucleotide sequence is shown in Fig. 1. That is, the recombinant vector of the present invention contains a nucleotide sequence which is the same or is functionally equal to the DNA having the nucleotide sequence shown in Fig. 3. A functionally equal nucleotide sequence means any DNA fragment which can be used in accordance with a substantially similar method to the present invention so as to obtain the substantially identical results, i.e. the production of an enzyme having the FMN reducing activity and the nitroreductase activity of luminous bacteria in a suitable host.

That is, the "functionally equal nucleotide sequence" means any DNA fragment which can encode a protein having the same amino acid sequence, even if the nucleotide sequence is different, or a DNA fragment which can code a protein having the FMN reducing activity and the nitroreductase activity, even if there is a slight difference in the amino acid sequence attributed to a slight difference in the nucleotide sequence. Typical examples are the nucleotide sequence of Fig. 3 and the nucleotide sequence of Fig. 1 into which a site-specific mutation may be introduced.

The nucleotide sequence in Fig. 1 will be described as follows:

Recently developed techniques make it possible to genetically endow a suitable microorganism with th ability to synthesize a protein or peptide normally made by another organism. The technique makes use of a fundamental relationship which exists in all living organisms between the genetic material, usually DNA, and the proteins synthesized by the organism. This relationship is such that the amino acid sequence of the protein is reflected in the nucleotide sequence of the DNA. There are one or more trinucleotide sequence groups specifically related to each of the twenty amino acids most commonly occurring in proteins. The specific relationship between each given trinucleotide sequence and its corresponding amino acid constitutes the genetic code. The genetic code is believed to be the same or similar for all living organisms. As a consequence, the amino acid sequence of every protein or peptide is reflected by a corresponding nucleotide sequence, according to a well understood relationship. Furthermore, this sequence of nucleotides can, in principle, be translated by any living organism.

The trinucleotides, termed codons, are presented as DNA trinucleotides, as they exist in the genetic material of a living organism. Expression of these codons in protein synthesis requires intermediate formation of messenger RNA (mRNA). The mRNA codons have the same sequences as the DNA codons, except that uracil is found in place of thymine. Complementary trinucleotide DNA sequences having opposite strand polarity are functionally equivalent to the codons, as is understood in the art. An important and well known feature of the genetic code is its redundancy, whereby, for most of the amino acids used to make proteins, more than on coding nucleotide triplet may be employed. Therefore, a number of different nucleotide sequences may cod for a given amino acid sequence. Such nucleotide sequences are considered functionally equivalent since they can result in the production of the same amino acid sequence in all organisms, although certain strains may translate some sequences more efficiently than they do others. Occasionally, a methylated variant of a purine or pyrimidine may be found in a given nucleotide sequence. Such methylations do not affect the coding relationship in any way.

The typical example is a plasmld vector into which the DNA fragment having the nucleotide sequence is introduced. As this kind of vector, there can be used pUC [C. Yanisch-Perron, J. Vieira and J. Messing, Gene, 33, p. 110-115 (1985)] and pIN III [Y. Masui, J. Coleman, M. Inouye, Experimental Manipulation of Gene Expression (ed. M. Inouye), Academic Press, p. 15 (1983)].

Fig. 7 shows a construction process of this recombinant vector (the expression vector).

That is, a vector pFR3 having a reductase gene is cleaved with restriction enzymes HincII and Stul to obtain a fragment including a coding region, and this fragment is then inserted into an Small site of a pUC8 plasmid DNA [Hanna Z., Fregeau C., Prefontaine G. and Brousseau R., Gene, p. 30247 (1984)] to construct a recombinant vector pFR5. Furthermor , this vector pFR5 is cleav d with a r striction enzyme EcoRI and then subjected to a Klenow treatment in the presence of dNTP. Afterward, the vector is recirculized using a T4 DNA ligase to central recombinant vector pFR7 (an expression vector). For the orientation of the thus construct deprod-

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uct, a restriction enzyme cleavage site is shown in an ampicillin resistant gene (Ampr).

Bact ria of the present invention contain a recombinant vector DNA having the nucleotide sequence shown in Fig. 1. The bacteria of the present invention are characterized by producing a protein having the flavin reducing activity and the nitroreductase activity.

A method for preparing the nzyme of the present invention comprises the steps of cultivating bacteria modified with a recombinant vector (an expression vector) containing a DNA whose nucleotide sequence is shown in Fig. 3, and then producing a protein containing an amino acid sequence shown in Fig. 4. Examples of the bacteria include Escherichia coli and Bacillus subtilis, and examples of a culture medium to be used include an LB culture medium and a YT culture medium.

A gene of the present invention is that which has been isolated for the first time encoding an enzyme having an FMN reducing activity and a nitroreductase activity. This gene can be used to produce a highly sensitive strain of bacteria to a mutagen or a carcinogen by the use of a suitable host such as <u>Escherichia coli</u>. Additionally, from this Escherichia coli, a reductase protein can also be prepared in large quantities.

By inserting this expression vector into a suitable host such as <u>Escherichia coli</u>, organisms or bacteria can be produced which express an enzyme having the FMN reducing activity and the nitroreductase activity of luminous bacteria. Furthermore, the reductase can also be prepared in large quantities by extraction from the organisms into which the gene is introduced. The organisms or microorganisms into which the gene is introduced have a high sensitivity to a mutagen or a carcinogen owing to the above-mentioned function, and thus they are useful as an indicator for detecting the mutagen or the carcinogen.

The reductase amplifies a luminous reaction of bacterial luciferase owing to the above-mentioned function. Thus, the reductase can be applied to many measuring methods and it is useful, for example, as a diagnosis drug or an inspection drug.

EXAMPLES

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Now, the isolation and identification of a gene which is important to the present invention will be described in reference to examples.

Example 1

[Identification of NAD(P)H:FMN reductase and determination of N-terminal amino acid sequence]

An NAD(P)H:FMN reductase sample (available from Boehringer Mannheim) was introduced into a "Sparose 12" gel filtration column (made by Pharmacya Co., Ltd.) to fractionate the sample. For each fraction, NADH and NADPH:FMN reducing activities were measured by a procedure described in Jablonski E. and DeLuca M., Biochemistry, 16, p. 2932 (1977), and analysis was then made in accordance with a procedure described in Laemmli, U.K. Nature, 277, p. 680 (1970) by means of sodium dodecyl sulfate-polyacrylamide gel electrophoresis.

As a result, it was clarified that the FMN reducing activity is directly proportional to the amount of a protein of 26 kDa (which is denoted by an arrow in Fig. 8).

After an SDS-PAGE analysis of this protein, it was transferred into a nylon membrane, and its amino acid sequence was determined in a usual manner by the use of a protein sequencer [made by Applied Biosystems Inc. (ABI)]. The results are set forth in Fig. 5. From these results, an N-terminal amino acid sequence having sequence numbers of 1 to 24 was confirmed.

Example 2

[Preparation of luminescent bacteria genomic library]

A photobacterium medium containing luminescent bacteria <u>Vibrio fischeri</u> (ATCC7744) was shaken at 26°C overnight to cultivate the bacteria. The bacteria were collected by means of centrifugal separation at 10000 rpm, and the resultant cell pellets were then dispersed in a Tris-HCl-EDTA buffer solution (hereinafter referred to as "a TE buffer"). After a lysozyme treatment at 37°C for 1 hour, sodium dodecyl sulfate (hereinafter abbreviated to "SDS") was added, followed by a proteinase K treatment at 50°C for 3 hours. Afterward, a phenol treatment was carried out thre times, followed by ethanol precipitation. After drying, the dried mat rial was dissolved in the TE buffer, and then subjected to the proteinase K treatment again. Afterward, the three cycles of the phenol treatment and then the ethanol precipitation were carried out to recover the genomic DNA. 10 units of a restriction enzyme Sau3AI were reacted with 50 µm of this genomic DNA at 37°C. Some parts of the reaction

mixtur were taken out at reaction times of 5, 10, 20, 30, 45, 60, 90 and 120 minutes, and afterward, EDTA (ethylen diaminetetraacetic acid) was added to the reaction system to bring the reaction to an ind. Each part of the DNA was subject d to agarose gel electrophoresis to confirm the degree of partial decomposition of the genomic DNA. The reaction solutions at the respective times were combined into one, followed by the ethanol precipitatin, to recover the DNA. Next, this DNA was dissolved in a small amount if the TE buffer, and then subjected to agarose gellectroph resist recover a fraction of 4 to 6 Kb by the use of a DE81 paper. The DNA fraction of 4 to 6 Kb was dissolved out of the DE81 paper with 1 M NaCl, subjected to the phenol treatment three times, and then precipitated with ethanol. The sample was dissolved in the TE buffer so as to be about 200 ng/µl. Afterward, the DNA fraction of 4 to 6 Kb was reacted with a pUC18 plasmid DNA (a plasmid vector), which was previously cleaved with a restriction enzyme BamH I and then treated with an alkaline phosphatase (an enzyme for catalyzing dephosphorization at the 5' terminal of the DNA), at 16°C overnight in the presence of a T4 DNA ligase (an enzyme for ligating DNA chains to each other or ligating the DNA and the 3'OH of an RNA or the 5'P terminal by a phosphodlester bond), whereby the DNA fraction was ligated to the plasmid. The resultant ligation reaction solution was transferred to JM109 Escherichia coli so as to perform transformation, and the thus obtained transductant represented a gene library.

[Preparation of synthetic oligonucleotide probe]

On the basis of the information of an amino acid sequence shown in Fig. 5, two probes of an oligonucleotide probe (FR-1) and an oligonucleotide probe (FR-2) were synthesized by means of a DNA synthesizer (made by ABI). Each synthetic probe was purified by the use of an OPC cartridge (made by ABI).

[Cloning of NAD(P)H:FMN reductase gene and analysis of its structure]

The gene library of Example 2 was screened in accordance with a colony hybridization method by the use of the FR-1 probe and the FR-2 probe. The FR-1 probe and the FR-2 probe were labelled at the 5' terminal with [y-32P]ATP and then used as labelled probes. After the titer of the gene library was measured, this g ne library was scattered on a nitrocellulose filter so as to be 200 colonies per plate. Cultivation was made at 37°C overnight, and two replicas were taken per filter. Each pair of two replicas was cultivated at 37°C and then used for hybridization. The filter was air-dried and then irradiated with ultraviolet rays (UV) to fix the DNA. Afterward, the filter was put in a hybridization solution {20 ml of a 6xSET buffer [20xSET buffer: 3 M of NaCl, 0.6 M of Tris-HCI (pH 8.0) and 0.04 M of EDTA], a 10xDehhardt's solution [(a solution containing 0.2% of each of serum albumin, polyvinylpyrrolidone and Ficoll), a 0.1% SDS and a salmon sperm DNA (thermally denatured, 50 μm/ml)], and it was then maintained at 68°C for 1 hour. Furthermore, the solution was replaced with a new one and then maintained for 1 hour, and a 32P-levelling probe was added, followed by hybridization at room temperature overnight. The solution was thrown away, and the filter was then washed with the 6xSET buffer, followed by shaking at 37°C for 20 minutes. After this operation was repeated twice, the filter was air-dried and then subjected to autoradiography. The filter was superposed upon a developed X-ray film, and the position of an ink marker was photographed on the film. Identification was made by aligning signals which were coincident with each other on the two films of the probe FR-1 and the probe FR-2 made from the one plate, and thus, five identified colonies (clones) were obtained.

[Preparation of recombinant vector]

For these five clones, a restriction analysis was carried out (Fig. 6), and as a result, it was apparent that three of these five clones were the same clones, which meant that three kinds of positive clones were prepared. Above all, a recombinant vector pFR3 (Fig. 7) having the smallest inserted DNA was used for the subsequent analysis.

50 [Structure determination of the gene and determination of amino acid sequence]

A Southern blotting analysis was made by the use of the FR-1 probe, and the region of an FMN reductase gene was determined in accordance with a dideoxynucleotide-enzyme method [Hattori M. and Sakaki Y., Anal, Biochem., 152, p. 232 (1986)], while reby a primary structure shown in Fig. 3 was elucidated. As a result, it was understood that the FMN reductas gene encoded a polypeptide of 24562 Da comprising 218 amino acids shown in Fig. 4, and this gene was about 30% homologous with a nitroreductase gene of Salmonella [Watanab M., Ishidate M. Jr and Nohmi T., Nucleic Acid Res., 18, p. 1059 (1990)].

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Example 3

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[Recombinant vector of NAD(P)H:FMN reductase gen , and construction of expression vector] (Fig. 7)

A recombinant v ctor pFR3 plasmid DNA was cleaved with a restriction enzyme Hinc II/Stu I and then treated at-80°C for 10 minutes. The thus treated DNA was then subjected to agarose gel electrophoresis to separate and recover a DNA fragment of about 1 Kb by the us of a DE81 paper. The DNA was diss lived out of the DE81 paper with 1 M NaCl, subjected to the phenol treatment three times, and then precipitated with ethanol. Next, the sample was dissolved in the TE buffer so as to be about 200 ng/μl. The above-mentioned DNA was reacted, at 16°C overnight in the presence of a T4DNA ligase, with a pUC8 plasmid DNA (a plasmid vector) which was previously cleaved with a restriction enzyme Sma I and then treated with an alkaline phosphatase, whereby the DNA was ligated to the plasmid. The resultant ligation reaction solution was transferred to JM109 Escherichia coli to perform transformation, and the Escherichia coli was selected and then cultivated overnight in a culture medium containing 5-bromo-4-chloro-3-indolyl-β-D-galactoside (Xgal) to form a white colony. This white colony was a transductant containing the plasmid into which the heterologous DNA was inserted.

A plasmid DNA was prepared from these transductants, and a restriction analysis was then carried out to obtain a strain containing a transformed vector pFR5. A plasmid DNA of the recombinant vector pFR5 was prepared, cleaved with EcoR I, subjected to a Klenow treatment, ligated with a T4DNA ligase, and then was transferred to D1210 Escherichia coli to perform transformation. Of the transductants, one in which an EcoR I cleavage site disappeared was selected. This was a recombinant vector (an expression vector) pFR7.

The recombinant vector pFR5 was constructed so as to express a peptide derived from a N-terminal β-galactosidase gene (lacZ) and a fused protein of the FMN reductase enzyme. The expression vector pFR7 was constructed so as to express lacZ and frameshift FMN reductase singly.

25 Example 4

[Preparation of Escherichia coli incorporated with NAD(P)H:FMN reductase gene]

Expression vectors pFR5 and pFR7 and a pUC8 plasmid DNA were transferred to D1210 <u>Escherichia coli</u> to perform transformation.

[Preparation of enzyme]

These transductants were incubated overnight, and 0.25 ml of the resultant incubation solution was transferred to an LB liquid (10 ml) culture medium containing ampicillin. After the culture medium was shaken at 37°C for 2 hours to cultivate the transductants, isopropyl-β-D(-)-thiolactopyranoside (hereinafter abbreviated to "IPTG") was added thereto so that a final concentration might be 1 mM, and the transductants were further cultivated for 3 hours. For the bacteria, an SDS-PAGE analysis was carried out to confirm the expression of a protein (this protein corresponds to the enzyme of the present invention).

The results are set forth in Fig. 4, but in the cases of the recombinant vectors pFR5 and pFR7, new bands appeared at 26 kDa which was the same size as in a commercial crude enzyme sample. In addition, in the case of the recombinant vector pFR5, a band appeared even at 29 kDa, and this vector was considered to be derived from a fused protein with lac Z.

1.5 ml of the incubation solution which was subjected to an IPTG induction treatment was centrifugally separated at 10000 rpm to remove a supernatant. The bacteria were dispersed ln 0.5 ml of a 50 mM potassium phosphate·1 mM dithiothreitol buffer, and then sonically disrupted by ultrasound. Centrifugal separation was further carried out at 4°C for 30 mlnutes at 12000 rpm, and the resultant supernatant was a cell extract.

For this cell extract, the following enzyme reducing activity was measured. The results are set forth in Tables 1, 2 and 3.

- (1) Flavin reducing activity: This was measured in accordance with a procedure described in Jablonski E and Deluca M., Biochemistry, 16, p. 2932 (1977).
- (2) Iron reducing activity: This was measured in accordance with a procedure described in Fontecave M., Eliasson R. and Reichard P., J. Biol. Chem., 262, p. 12325-12331 (1987).
- (3) Nitroreductase activity: This was measured in accordance with a procedure described in Watanabe M., Ishidate M. Jr. and Nohmi T., Mutation Research, 216, p. 211-220 (1989).

Protein amounts in the respective tables were determined in accordance with a Bradford method by thus of a protein assary kit made by Bio-RAD [Bradford M. M., Anal. Biochem., 72, p. 248-254 (1976)].

	1	1	1		L
5			(-)	80 20 00	110 110 50 0
10		ty n)	Riboflavin (+)	2120 2700 10 10	870 870 0 10
15		Flavin Reducing Activity (nmol/min/mq protein)	(-)	240 340 40 10	70 140 10 30
20		lavin Reducir (nmol/min/mg	FAD (+)	3980 7410 50 40	1680 28 4 0 0 10
25	le 1	Flavi			·
	Table		FMN (-)	430 660 30 30	190 230 30 50
30			÷	6330 11810 20 40	2300 4430 50 20
35			ų (c	.210 .210 1210 0	1210 1210 1210 0
40			Strain (IPTG)	pFR5/D1210 pFR7/D1210 pUC13/D1210 D1210	pFR5/D1210 pFR7/D1210 pUC13/D1210 D1210
45					
50				NADH	

5			Riboflavin (+)	2.9 4.2 0.0	0000
10		Activity protein)	Ribof	30.1 42.8 0.1	9.4 14.0 0.2 0.4
15		Se	FAD (-)	3.6 1.7 1.8	4.0 0.0 0.0
20		Iron Reducta (nmol/min/my	F (+)	39.4 59.5 1.6 1.3	11.8 23.6 0.1 0.1
25	Table 2		FMN (-)	4.5 6.6 0.1	0.0 0.3 4.0
30			F)	44.9 68.3 0.4	12.6 25.8 0.6 0.6
35			in G)	1210 1210 1210 1210 .0	1210 1210 1210 1210 .0
40			Strain (IPTG)	pFR5/D1210 pFR7/D1210 pUC13/D1210 D1210	pFR5/D1210 pFR7/D1210 pUC13/D1210 D1210
45				NADH	NADPH
50	l			Ż	ž l

Tabl 3

		Strain (IPTG)	Nitroreductase Activity (nmo	l/min/mg protein) Nitrofurazone
5			(+)	(-)
	NADPH	pFR5/D1210	24.8	10.1
		pFR7/D1210	40.9	10.4
10		pUC13/D1210	4.7	5.2
		D1210	6.8	5.5

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Comparing the activities in these tables, the activities of pFR5 and pFR7 are about 1 to 3 orders higher than those of pUC13 which is a negative control. This gene could therefore be identified as a gene encoding an enzyme protein having a flavin reducing activity and a nitroreductase activity.

Claims

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- 1. An isolated and purified gene containing a nucleotide sequence shown in Fig. 1 and encoding an enzyme having a flavin reducing activity and a nitroreductase activity.
- 2. An isolated and purified gene containing a nucleotide sequence shown in Fig. 2 and encoding an enzyme having a flavin reducing activity and a nitroreductase activity.

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- An isolated and purified gene containing a nucleotide sequence shown in Fig. 3 and encoding an enzyme having a flavin reducing activity and a nitroreductase activity.
- An enzyme containing an amino acid sequence shown in Fig. 4 and having a flavin reducing activity and a nitroreductase activity.
- 5. A recombinant vector containing a DNA whose nucleotide sequence is shown in Fig. 1.
- The recombinant vector according to Claim 5 wherein the nucleotide sequence shown in Fig. 1 is insert d into a plasmid vector.

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- 7. A bacterial host containing a recombinant vector which vector contains a DNA whose nucleotide sequence is shown in Fig. 1.
- A method for preparing an enzyme containing an amino acid sequence shown in Fig. 4 which comprises 40 a step of cultivating bacteria modified with a recombinant vector containing a DNA whose nucleotide sequence is shown in Fig. 1.

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ATG	VCT	CAK	CCL	ATM	MTA	CAK	GAK	XTY	GAJ	AAK	WGZ	TAK	ACL	QRS	AAJ	48
Met	Thr	His	Pro	He	Πe	His	Λsp	Leu	Glu	Asn	Λrg	Tyr	Thr	Ser	Lys	
1				5					10					15		
														GTL		96
Lys	Tyr	Asp		Ser	Lys	Lys	Val		GIn	GIn	Λsp	Leu		Val	Leu	
VጥV	CAT	CCI	. 20	W(*7	VTV	ODC	CCI	25	ODC	A T U	A A 1/	ADC	30	CCL	ጥር ር	144
														Pro		144
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۸۸J	TTK		GTL	ΛTM	GA.f	QRS		GCA	GCL	ΛAJ	CAJ		ATG	CAK	GAK	192
														His		
	50					55	_				60	-			_	
QRS	TTK	GCL	$\Lambda\Lambda K$	ATG	CAK	CAJ	TTK	AAK	CAJ	CCL	CAK	ATM	۸۸J	GCL	TGK	240
Ser	Phe	Ala	Asn	Met	His	GIn	Phe	Asn	Gln	Pro	llis	He	Lys	Ala	Cys	
65					70					75					80	
QRS	CAK	GTG	ATM	XTY	TTK	GCL	AAK	AAJ	XTY	QRS	TAK	ACL	WGZ	GAK	GAK	288
Ser	llis	Val	He		Phe	Ma	Asn	Lys		Ser	Tyr	Thr	Arg	Лsp	Лsp	•
				85					90					95		
														VCL		336
Tyr	Asp	Val		Leu	Ser	Lys	Ala		Λla	Asp	Lys	Λrg		Thr	Glu	
	~		100					105					110			
														TTG		384
61U	Gin		GIB	Ala	Ala	l'he		Ser	Phe	Lys	Phe		Glu	Leu	Λsn	
TOM	OAK	115	4 4 17	001	04.7	041	120					125				
														GCL		432
Cys		GIU	Asn	Gly	GIU		Lys	Ala	Trp	lhr		Pro	Gin	Ala	Tyr	
VTV	130	VTV	CCI	AAV	CCI	135	CAV	101	VTV		140	VANU		1.771	017	400
														ATM		480
145	nia	Leu	Gly	ASII	150	Leu	1115	mr	ren	155	Arg	reu	Asn	He	_	
	ACI	ACI	ATC	CAT		ΑТИ	CAY	CCI	CAI		TTI	UDG	CAT	ATM	160	528
														Ile		320
.,01		1111	m C L	160	OI,	110	пор	170	170	Leu	Leu	net	(71 ti	175	THE	
GCI.	GAK	GAI	хтч		GGL.	TAK	€: A I	TCK		GTI	CCI	XTV	cer	MTA	(20)	576
														He		010
	, F		180	,	,	• , •	~ · · · ·	185				Den	190		O, y	
TAK	CAK	CAK		QRS	GA."	GAK	TAK		GCL	QRS	TTG	CCL		QRS	WGZ	624
														Ser		021
-		195				•	200					205	_ , _		0	
۸۸J	CCL		GAJ	GCL	GTI.	ATM		ATM	XTY	TJJ		, •				657
						He										
	210					215										

								CTT								48
Met	Thr	His	Pro	He	He	llis	Λsp	Leu		Λsn	Arg	Tyr	Thr		Lys	
1				5					10					15		
								TC T								96
Lys	Tyr	Asp		Ser	Lys	Lys	Val	Ser	Gln	Glu	Λsp	Leu		Val	Leu	
			20			mo#		25	mo i	4 mm		T C.	30	CCT	TOO	144
								TCT								144
Leu	Glu		Leu	Arg	ren	Ser		Ser	Ser	116	ASII	5er 45	GIN	rro	ITP	
	ጥ ፓር	35	ርጥጥ	A TT	CAL	TCC	40	GCA	ecc	AAC	CAA		ATC	CAT	CAT	192
								Ala								102
Lys	50	116	181	116	OTU	55	nsp	nia	nia	Lys	60	оту	MCC	игэ	лор	
TCC		CCA	AAT	ATC	CAT		т т т	AAT	CAA	сст		ATC	AAA	GCG	TGT	240
								Asn								
65	1110	nia	non	мсь	7()	0111	1 110	.1011	• • • • • • • • • • • • • • • • • • • •	75		1.0	, .		80	
	CAT	GTG	Λ T T	TTA		GCA	AAT	AAG	CTT		TAT	ACA	CGA	GAT		288
								Lys								
				85					90					95		
TAT	GAT	GTG	GTT	TTA	TCT	$\Lambda A A$	GCG	GTT	GCT	GAC	AAG	CGT	ΛTT	ACT	GAA	336
Tyr	Asp	Val	Val	Leu	Ser	Lys	Ala	Val	Ala	Asp	Lys	Arg	He	Thr	Glu	
			100					105					110			
								TCG								384
Glu	Gln	Lys	Glu	Ala	Ala	Phe	Ma	Ser	Phe	Lys	Phe			Leu	Asn	
		115					120		_			125				
								GCA								432
Cys	_		Asn	Gly	Glu		Lys	Ala	Trp	Thr	4	Pro	Gin	Ala	Tyr	
	130		0.05		0.00	135	0.45		TT 4	COT	140	CTC.	110	4 TT	CAC	400
								ACA								480
	Ala	Leu	ыу	ASN			nıs	Thr	Leu	155	MI g	Leu	ASII	116	160	
145	ACA	ACA	ATC	CAA	150 ccc		(2 A T	CCT	CAA		ттс	ACT	CAA	АТТ		528
								Pro								020
OCI	1111	1111	met	160		110	1131	1117	170	DCu	Deu	.,,,	oru	175		
GCT	GAT	GAA	TTA			TAT	GAA	TGT		GTT	GCT	ТТА	GCC			576
								Cys								
	•		180	,	•	•		185					190			
TAT	CAT	CAT		AGC	GAA	GAT	TAT	ΛΑΤ	GCC	TCT	TTG	CCT	AAG	TCT	CGT	624
															Arg	
-		195					200					205				
AAG	GCA	TTT	GAA	GCA	GTA	ATT	VCC	ATC	CTT	TAG						657
Lys	Ala	Phe	Glu	Ala	¥a1	He	Thr	He	Leu	***						
	210					215										

													AT(Me	G ACC	TGTGC G CAT His	60 117
				GAT Asp									AAA	TAT		165
	TCA			GTA Val		$C\Lambda\Lambda$					GTT					213
CTG				GCT Ala 40	TCT					CAG					ATT	261
				GAT Asp					GGT					TTT		309
			CAG	TTT Phe				CAC					TCT			357
		TTT		AAT Asn			TCG					GAT				405
	TTA			GCG Ala		GCT					ACT				Lys	453
GAA				GCT Ala 120	TCG				Val	GAA				Asp		501
AAT Asn	GGT Gly	GAG Glu	llis	AAA Lys	GCA Ala	TGG Trp	ACT Thr	Lys	125 CCT Pro	CAA Gln	GCT Ala	TAT Tyr	Leu	130 GCT Ala	CTT Leu	549
GGT Gly	AAT Asn	GCT Ala 150	135 CTG Leu	CAT His	ACA Thr	TTA Leu	GCT Ala 155	140 AGA Arg	CTG Leu	AAC Asn	ATT He	Asp	145 TCA Ser	ΛCΛ Thr	ACA Thr	597
		GGC		GAT Asp		Glu	TTA				He					645
TTA Leu 180	AAA	GGG Gly	TAT Tyr	GAA Glu	TGT Cys 185	170 CAT Ilis	GTT Val	GCT Ala	TTA Leu	GCC Ala 190	175 ATT Ile	GGT Gly	TAT Tyr	CAT His	His	693
CCA				TAT Tyr	AAT				Pro	AAG				Λla		741
				ACC Thr					205					210		762
TAG! ***	\TTC'	TTA .	ATGT:	TTGAG	GA TO	GAAGA	AAAA	; cc	AGCG	TTT	AGC	rgtgg	CTT :	rgtt:	TGTGCA	822
AAA				GGCG CAAT									TTT (CTTT	глстст	882 929

Met 1	Thr	His	Pro	Ile 5	He	His	Asp	Leu	Glü 10	Asn	Arg	Tyr	Thr	Ser 15	Lys
Lys	Tyr	Asp	Pro 20	Ser	Lys	Lys	Val	Ser 25	Gln	Glu	Λsp	Leu	Ala 30	Val	Leu
Leu	Glu	Ala 35	Leu	Arg	Leu	Ser	Ala 40	Ser	Ser	Ilc	Asn	Ser 45	GIn	Pro	Trp
Lys	Phe 50	He	Val	He	Glu	Ser 55	Λsp	Ala	Ala	Lys	G1 n 60	Gly	Met	llis	Asp
Ser 65	Phe	Ala	Asn	Met	llis 70	G1n	Phe	Asn	G1n	Pro 75	lłis	Ile	Lys	Лlа	Cys 80
Ser	His	Val	He	Leu 85	Phe	Ala	Asn	Lys	Leu 90	Ser	Tyr	Thr	Arg	Asp 95	Asp
Tyr	Лsp	Val	Val 100	Leu	Ser	Lys	Λla	Val 105	Ala	Asp	Lys	Arg	11e 110	Thr	Glu
Glu	Gln	Lys 115	Glu	Ala	Ala	Phe	Ala 150	Ser	Phe	Lys	Phe	Val 125	Glu	Leu	Asn
Cys	Asp 130	Glu	Asn	Gly	Glu	llis 135	Lys	Ala	Trp	Thr	Lys 140	Pro	Gln	Ala	Tyr
Leu 145	Ala	Leu	Gly	Asn	Ala 150	Leu	llis	Thr	Leu	Ala 155	Arg	Leu	Asn	He	Λsp 160
Ser	Thr	Thr	Met	Glu 165	Gly	He	Asp	Pro	Glu 170	Leu	Leu	Ser	Glu	He 175	Phe
Λla	Asp	Glu	Leu 180	Lys	Gly	Tyr	Glu	Cys 185	llis	Val	Ala	Leu	Ala 190	He	Gly
Tyr	His	His 195	Pro	Ser	Glu	Asp	Tyr 200	Λsn	Ala	Ser	Leu	Pro 205	Lys	Ser	Arg
Lys	Ala 210	Phe	Glu	Ala	Val	I1c 215	Thr	Ile	Leu						

Fig.6

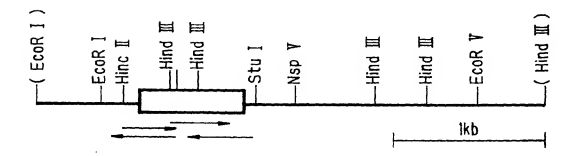


Fig.7

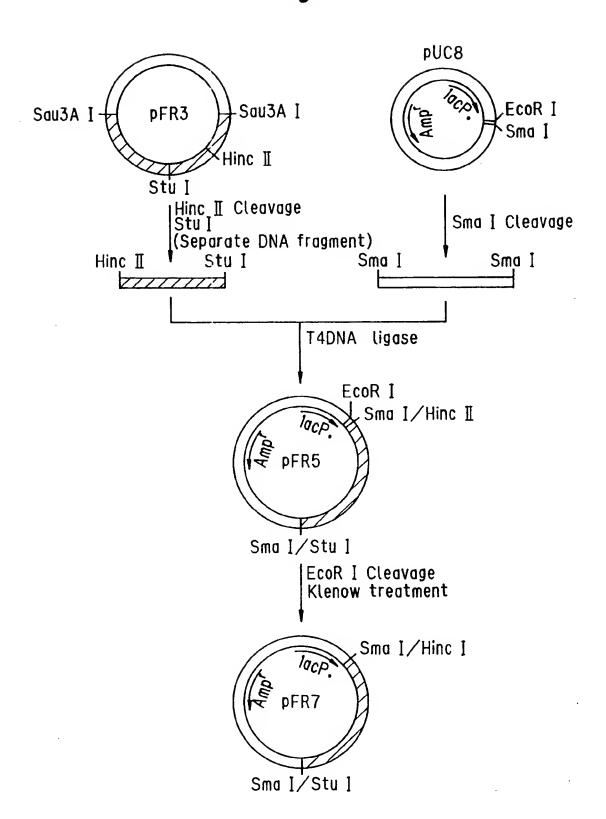
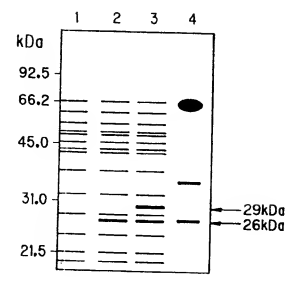


Fig.8





EUROPEAN SEARCH REPORT

Application Number

EP 92 31 1476 PAGE1

				LVGET
	DOCUMENTS CONSI	DERED TO BE RELEVAN	T	
Category	Citation of document with in of relevant pas	dication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.5)
D,Y	pages 3673 - 3679 G. SPYROU ET AL. ¹Ch	une 1991, BALTIMORE, US naracterization of the ne (fre) of Escherichia on of a plasmid for ne enzyme ^t		C12N15/53 C12N9/02 //(C12N9/02, C12R1:63)
	JOURNAL OF BIOLUMINE CHEMILUMINISCENCE vol. 5, no. 3, 1990, pages 187 - 192 J.T. LAVI ET AL. 'Afbacterial luciferase oxidoreductases by Fanalytical applicati* the whole document	ffinity purification of e and NAD(P)H:FMN FMN-Sepharose for ions'	1-8	
	US pages 4119 - 4125 C. BRYANT ET AL. 'Pu characterization of	March 1991, BALTIMORE, Prification and an oxygen-insensitive Use from Enterobacter	4	TECHNICAL FIELDS SEARCHED (Int. Cl.5)
	US pages 2932 - 2936 E. JABLONSKI ET AL. properties of the NA	ADH and NADH specific from Beneckea harveyi' , * 	1-8	
	Place of search	Date of completion of the search		Examiner
В	BERLIN	16 MARCH 1993		JULIA P.
X : parti Y : parti docu A : tech O : non-	CATEGORY OF CITED DOCUMEN icularly relevant if taken alone icularly relevant if combined with anot meet of the same category mological background written disclosure mediate document	E : earlier patent doc after the filing da	ument, but publite in the application or other reasons	is hed an, ar

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EUROPEAN SEARCH REPORT

Application Number

EP 92 31 1476 PAGE2

Category	Citation of document with it of relevant pa	ndication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CL5)
Y	JOURNAL OF BIOLOGIC vol. 266, no. 7, 5 US pages 4126 - 4130 C. BRYANT ET AL. 'C sequence, and expre nitroreductase gene cloacae' * the whole documen	March 1991, BALTIMORE loning, nucleotide ssion of the from Enterobacter	1-8	
D,A	the detection of mu construction of nitroreductase-over	'A sensitive method 1 tagenic nitroarenes: producing derivatives murium strains TA98 a		
A		ial NAD(P)H: flavin	4	TECHNICAL FIELDS SEARCHED (Int. Cl.5)
D,A	NUCLEIC ACIDS RESEA vol. 18, no. 4, 199 page 1059 M. WATANABE ET AL. Salmonella typhimur gene ¹ * the whole documen	O, LONDON, GB 'Nucleotide sequence ium nitroreductase	1-4 of	
	The present search report has b	een drawn up for all claims Date of consistion of the surre		Permiser
1	BERLIN	16 MARCH 1993		JULIA P.
Y : par doc	CATEGORY OF CITED DOCUME ticularly relevant if takes alone ticularly relevant if combined with an ament of the same category shoological background	E : earlier pate after the fit other D : document o	rinciple underlying the new document, but put ling date free in the application ited for other reasons	olished on, or